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Title: Opposites attract: MHC-associated mate choice in a polygynous primate

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Short running title: MHC-associated mate choice in a primate

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Abstract

We investigated reproduction in a semi-free-ranging population of a polygynous primate, the mandrill, in relation to genetic relatedness and male genetic characteristics, using neutral microsatellite and MHC genotyping. We compared genetic characteristics of the sire and genetic dissimilarity to the mother with all other potential sires present at the conception of each offspring (193 offspring for microsatellite genetics, 180 for MHC). The probability that a given male sired increased as pedigree relatedness with the mother decreased, and overall genetic dissimilarity and MHC dissimilarity with the mother increased. Reproductive success also increased with male microsatellite heterozygosity and MHC diversity. These effects were apparent despite the strong influence of dominance rank on male reproductive success. The closed nature of our study population is comparable to human populations for which MHC-associated mate choice has been reported, suggesting that such mate choice may be especially important in relatively isolated populations with little migration to introduce genetic variation.

Keywords: major histocompatibility complex, disassortative mating, good genes, heterozygosity; sexual selection

INTRODUCTION

Mate choice, particularly female choice, has been the focus of extensive research over the past two decades (Andersson & Simmons, 2006). Where there is little or no direct benefit of mate choice to an individual or its offspring, females may choose for genetic benefits that will be inherited by their offspring (choice for 'good genes'). These indirect benefits may include increased offspring attractiveness (Fisher, 1958) or other heritable qualities (Zahavi, 1975) such as immunocompetence and parasite resistance (Hamilton & Zuk, 1982; Folstad & Karter, 1992). Adaptive complementarity may also be an important factor in mate selection (Trivers, 1972; Zeh & Zeh, 1996) since offspring born to closely related parents often show reduced fitness (inbreeding depression) (Keller & Waller, 2002). Estimators of genetic diversity are correlated with a range of fitness components, including survival, disease susceptibility, and reproductive success (review in Hansson & Westerberg, 2002). Females should therefore benefit by mating preferentially with genetically different males, thereby increasing the heterozygosity of their progeny. However, choice for genetically dissimilar mates may trade-off against the loss of locally adaptive gene complexes, leading to choice for some optimal level of dissimilarity (Bateson, 1983).

The Major Histocompatibility Complex (MHC) is among the best candidates for the genetic basis of mate choice in vertebrates (Jordan & Bruford, 1998; Penn & Potts, 1999). The MHC is a multigene family encoding cell-surface glycoproteins (MHC molecules) that play a critical role in the immune system by recognising foreign peptides, presenting them to specialist immune cells and initiating the appropriate immune response (Klein, 1986). Expressed loci are highly polymorphic and this diversity is selectively maintained, at least in part, via two mechanisms of pathogen-mediated selection: heterozygote advantage and frequency-dependent selection (Apanius *et al.*, 1997; Sommer, 2005). In the former mechanism, heterozygote individuals are able to resist a wider range of pathogens, rendering them fitter than less diverse individuals (Doherty & Zinkernagel, 1975). In the latter, a particular allele is beneficial when rare, but disadvantageous when common, because natural selection favours parasites that can evade the MHC-dependent immunity of the most common host genotypes, decreasing the fitness of individuals possessing common alleles. Rare

alleles are thus favoured, because they escape recognition by the MHC-dependent immune system, until they increase in frequency and parasites evolve to evade them, in a co-evolutionary arms race (Penn & Potts, 1999).

MHC-based mate choice may favour individuals that possess particular MHC alleles, those with diverse MHC genotypes, or those with MHC genotypes that are dissimilar to the chooser (review in Penn & Potts, 1999; Penn, 2002). Choice for particular beneficial alleles may provide offspring with resistance to particular parasites (Penn & Potts, 1999). Choice for an MHC-diverse mate may be advantageous because heterozygotes possess more rare alleles than homozygotes, which can be inherited by offspring, and because an MHC-diverse mate is less likely to share alleles with the chooser, leading to MHC-diverse offspring, that are able to resist a broader range of pathogens (Apanius *et al.*, 1997; Fromhage *et al.*, 2009). Finally, mate choice for MHC dissimilarity (disassortative mating) may provide several, non-exclusive, fitness benefits: preventing inbreeding and increasing genome-wide genetic diversity (Brown & Eklund, 1994); increasing the ability of offspring to resist pathogens through either heterozygote advantage (Zuk, 1990) or the production of offspring that are dissimilar to the parents (Penn & Potts, 1999); or giving offspring an optimal number of MHC alleles for parasite resistance ('allele counting') (Nowak *et al.*, 1992; Reusch *et al.*, 2001; Wegner *et al.*, 2003; Forsberg *et al.*, 2007) (but see Borghans *et al.*, 2003).

Support for MHC-based mate choice hypotheses was first obtained from studies of laboratory mice (Yamazaki *et al.*, 1976). More recently, evidence that the MHC influences mate choice has come from studies of fish, birds and mammals (review in Piertney & Oliver, 2006). However, few studies have examined MHC-associated mate choice in non-model species living in natural, or semi-natural, populations (Piertney & Oliver, 2006). Of the studies that exist, some have found evidence for choice for MHC-dissimilar mates (Landry *et al.*, 2001), some that females choose males to achieve an intermediate, and optimally resistant, level of MHC diversity in their offspring (Milinski *et al.*, 2005; Bonneaud *et al.*, 2006), and still other studies found no influence of the MHC on mate choice at all (Paterson & Pemberton, 1997; Ekblom *et al.*, 2004; Westerdahl, 2004). These studies suggest that MHC-associated mate choice may occur in some species, but not in others, and that the exact strategies employed may differ between

species (Piertney & Oliver, 2006). Furthermore, most studies of MHC-associated mate
choice have failed to include expression analyses, and it remains to be seen whether the
MHC sequences studied actually produce functional molecules for pathogen resistance
(Knapp, 2007).

The role of the MHC in human mate choice is particularly controversial. Initial studies
suggested that MHC dissimilarity plays a role in human mate choice (Ober et al., 1997),
and experiments suggest that this phenomenon may be mediated via odour (Wedekind
et al., 1995; Wedekind & Furi, 1997; Jacob *et al.*, 2002). However, other studies found no
influence of MHC dissimilarity on human mate choice (Hedrick & Loeschcke, 1996;
Hedrick & Black, 1997; Ihara *et al.*, 2000; Chaix *et al.*, 2008). This controversy extends to
non-human primates. A study of group-living rhesus macaques (*Macaca mulatta*) found
no evidence of mate choice for MHC-dissimilarity, although MHC-heterozygous males
enjoy increased reproductive success (Sauermaun et al., 2001). However, female choice
for both MHC dissimilarity and within-male MHC diversity and, as well as for males with
higher genome-wide heterozygosity, has been reported for socially monogamous fat-
tailed dwarf lemurs (*Cheirogaleus medius*) (Schwensow et al., 2007a) and solitary
foraging grey mouse lemurs (*Microcebus murinus*) (Schwensow et al., 2008).

We investigated the influence of MHC genotype on patterns of reproduction in the
mandrill (*Mandrillus sphinx*, Cercopithecinae). Mandrills live in large multi-male, multi-
female groups (Abernethy et al., 2002), and are moderately seasonal breeders (Setchell
& Wickings, 2004). The potential for male-male contest to monopolise access to
individual receptive females is thus high, and mandrills have a polygynous mating
system, with strong sexual dimorphism (Setchell et al., 2001) and high reproductive
skew in favour of the alpha male (Charpentier et al., 2005a). Nevertheless, female
mandrills are able to mate with multiple males during a single receptive period
(Setchell, unpublished observations), and express precopulatory mate choice (Setchell,
2005). Female mandrills gain little in the way of direct benefits from males and female
choice is likely, therefore, to be driven by the potential indirect (genetic) benefits that a
sire may provide. Both inbreeding and the reduction of genome-wide heterozygosity
have deleterious consequences for individual fitness (Charpentier *et al.*, 2005b; 2006)
meaning that mate choice for non-relatives and/or genetically complementary

individuals would produce more heterozygous, fitter progeny. However, the relatively tight control that dominant males appear to have over both mating and paternity may reduce the ability of females to reproduce with non-dominant males of their choice, as proposed for Soay sheep (Paterson & Pemberton, 1997). We also test for the possibility that within-male MHC diversity, or the possession of particular MHC types, confer a reproductive advantage on males, via either superior competitive ability (intra-sexual selection), or via female choice for such males.

We genotyped a large population of mandrills for a highly variable group of MHC class II loci known as MHC-DRB genes. These genes encode proteins that are directly involved in immune response and are under strong positive selection pressure with the peptide binding region containing significantly more non-synonymous than synonymous changes (Abbott et al 2006), suggesting that this area of the genome is under balancing selection. We also demonstrated that many of the MHC sequences we identified via genomic DNA analysis are expressed. Next, we compared genetic and demographic characteristics of the sire of each individual offspring with all the potential sires available when the individual was conceived, to address four specific questions: (1) Do mandrills choose genetically dissimilar mates to avoid inbreeding? (2) Do mandrills mate disassortively based on MHC genotype? (3) Do males with greater overall genetic diversity, or greater within-male MHC diversity, experience greater reproductive success? (4) Do specific MHC genotypes influence male reproductive success? We found that the probability that a given male sired increased as pedigree relatedness decreased, and overall genetic dissimilarity and MHC dissimilarity with the mother increased. Reproductive success also increased with male microsatellite heterozygosity and within-male MHC diversity. These effects were apparent despite the strong influence of dominance rank on male reproductive success.

METHODS

Study population

We studied a large, semi-free-ranging population of mandrills, at the Centre International de Recherches Médicales, Franceville (CIRMF), Gabon, established in 1983/4, when 15 wild founder were released into a 6.5 ha naturally rain-forested

enclosure (see Setchell et al. 2005 for details of the colony). The date of birth is recorded for all individuals born into the colony, while the age of founder animals was approximated using dental estimates when the animals arrived at CIRMF and their previous history. Daily observations are made of female reproductive status, births, injuries and disappearances. Male rank is determined on the basis of avoidance behaviours; the identity of the top-ranking (alpha) male is unambiguous. Paternity skew is concentrated in alpha males, and beta males do not sire more offspring than other subordinate males (Setchell *et al.*, 2005a), so we limit comparisons to alpha vs. non-alpha males.

Group sizes ranged from 15 in 1983/4 to a maximum of 104 animals in 2002, corresponding to smaller groups observed in the wild (Rogers et al., 1996). How the situation in the colony relates to wild mandrills is currently unknown, but it seems likely that the restricted conditions of the CIRMF colony represent an extreme, but not totally un-natural, situation (Setchell *et al.*, 2005b).

Microsatellite genotyping and paternity

We extracted DNA for genetic analyses from blood samples obtained during annual captures of the colony. We genotyped up to ten microsatellite loci for 14 founder animals and 205 offspring born into the colony between 1983 and 2002. We obtained an accurate assignment of paternity for 193 (94%) of 205 offspring (for details of methods and paternity assignment criteria, see Charpentier et al., 2005a).

MHC genotyping

We conducted MHC-DRB genotyping for 155 of the study population (insufficient DNA was available for the remaining individuals). We PCR amplified MHC-DRB sequences using primers known to amplify all MHC-DRB sequences in species ranging from humans to New World monkeys and analysed products using denaturing gradient gel electrophoresis (DGGE) and direct sequencing (Abbott et al., 2006). We amplified DNA samples from each individual multiple times and repeated all genotyping experiments to ensure that any sequence found in one individual would also be detected in all other individuals in the population.

The MHC-DRB region in Old World primates frequently experiences expansion and contraction through gene duplication and deletion, respectively (Sliereendregt *et al.*, 1994). Due to the extensive variation in DRB haplotype composition, individuals possess different numbers and types of DRB genes on each haplotype. We therefore focus on the number of different sequences possessed by an individual as a measure of MHC diversity, without making any assumptions about the number of loci involved (see also Málaga-Trillo *et al.*, 1998; Aeschlimann *et al.*, 2003; Ekblom *et al.*, 2004; 2008).

To determine whether the mandrill MHC sequences produce functional molecules for pathogen resistance we examined patterns of expression using cDNA analysis for a subset of seven mandrills chosen to represent all known Masp-DRB loci and lineages. We calculated the number of amino-acid differences between each pair of MHC sequences as an estimate of genetic dissimilarity (Landry *et al.*, 2001), because MHC sequences may differ in nucleotide composition, but be functionally similar in terms of immune defence if the protein they encode binds the same peptides (Rammensee, 1995; Sidney *et al.*, 1995). We also used MHC-DRB sequences to determine MHC-DRB supertypes. These are groups of MHC-DRB sequences that share peptide-binding motifs and are therefore functionally similar (Doytchinova & Flower, 2005), and have been shown to be biologically relevant in studies of both human and non-human primates (Southwood *et al.*, 1998; Trachtenberg *et al.*, 2003; Schwensow *et al.*, 2007b). We identified variable amino acid positions, presumed to represent the peptide binding region, using phylogenetic analysis of MHC sequences in MEGA 4 (Tamura *et al.*, 2007). We then used PAML 4 (Yang, 2007) to identify positively selected sites (PSS). Finally, we identified supertypes by analysing the chemical specificities of these PSS in Genesis version 1.7.2 (Sturn *et al.*, 2002), following Doytchinova and Flower (2005).

Relatedness and reproduction:

To determine whether reproduction was biased towards unrelated partners we estimated the overall genetic similarity between the genotypes of two individuals as:

R_{ped} A relatedness coefficient calculated using the colony pedigree (R_{ped} in mother-son and father-daughter pairs is 0.5, full-siblings 0.5, half-siblings 0.25, etc.)

240 R_{QG} Microsatellite allele-sharing, calculated as the Queller-Goodnight index
(Queller & Goodnight, 1989) using RELATEDNESS (Version 5.0.8;
242 available from www.gsoftnet.us/GSoft).
We also classified R_{ped} as >0.25 (i.e. father/daughter dyads and half-siblings) and <0.25
244 for some analyses ($R_{<0.25}$).

246 ***MHC-dissassortative mating***

To determine whether reproduction was biased towards partners with dissimilar MHC
248 genotypes, we calculated three measures of MHC dissimilarity for each potentially
reproductive dyad:

250 MHC_{diff} The number of MHC sequences that differed between the male and
female. This was highly and significantly correlated with the number of
252 MHC sequences shared and the number of MHC sequences unique to the
male so we report only results for MHC_{diff} .

254 AA_{diff} Amino acid sequence dissimilarity, calculated as the mean number of
pairwise amino acid differences between the sequences of the dyad.

256 S_{diff} The number of MHC supertypes that differed between the male and
female.

258

Male genotype and reproduction

260 To determine whether reproduction was biased towards males that were more
genetically diverse, possessed higher MHC diversity, or possessed particular MHC
262 supertypes, we described the genotype of a potential sire as follows:

IR_{male} Internal Relatedness (IR, Amos et al., 2001). The more an individual is
264 genetically diverse, the more IR is negative. While measures of
heterozygosity based on small number of neutral markers may not
266 accurately reflect genome-wide heterozygosity (Balloux *et al.*, 2004; Slate
et al., 2004), we have previously shown that our measure of IR is a good
268 measure of genome-wide inbreeding in this population (Charpentier et al.,
2005b).

270 MHC_{male} Number of MHC sequences possessed.

AA_{male} MHC sequence diversity, calculated as the mean number of amino acid
272 differences between all MHC sequences.

Smale Number of supertypes possessed.

274 S1 to S13 The presence/absence of individual MHC supertypes.

276 ***Statistical analyses***

We conducted statistical analyses at the level of the individual offspring, asking the
278 following question ‘based on the potential sires available, their genetic similarity to the
female, and their individual genetic characteristics, which male sired the offspring?’
280 Potential sires were any adolescent (4-9 yrs) or adult male (>9yrs, Setchell *et al.*, 2006)
present in the group at the time that the mother conceived. Our microsatellite dataset
282 contained 193 offspring, 51 potential sires (1-113 potential offspring per sire, mean
46±5), 17 actual sires, (1-42 true offspring per sire, mean 11.4±3), and 42 mothers (1-
284 15 offspring per female, mean 4.6±0.7). The MHC dataset contained 180 offspring, 40
potential sires (1-109 potential offspring per sire, mean 45±5), 15 actual sires (1-42
286 true offspring per sire, mean 12±3.2), and 34 mothers (1-15 offspring each, mean
5.3±0.8). The same potential sires and mothers appeared several times in our dataset.
288 However, the number and identity of potential sires available differed for each offspring
born to an individual female because the males available as female mandrills conceive
290 approximately one infant per year (Setchell *et al.*, 2005b) and potential sires differed
across breeding seasons. Thus, while a potentially reproducing dyad could appear more
292 than once in the data-set, the set of alternative potential sires (i.e. the ‘choice’ of sire
available) for a given female was different for each of her offspring.

294

Our dependent variable (‘decision’) took the value 1 when a given male was identified
296 as the sire of the offspring; and 0 for all other potential sires present in the group at the
time of conception. This decision variable does not follow a binomial distribution
298 because only one potential sire scored ‘1’ for each offspring, while all other scored ‘0’.
To resolve this problem we used conditional logit regression models (multinomial
300 discrete choice: MDC procedure, type=clogit, SAS version 9) to investigate the influence
of different variables on the probability that a given male sired. The MDC procedure
302 analyses models where the choice set consists of multiple alternatives, in this case
multiple potential sires for each offspring. The model takes into account the number of
304 sires available to sire that particular individual. It also takes into account the identity of
the males, and therefore their repetition throughout the dataset. However, the model

does not consider the fact that some mothers contributed more than one offspring to the dataset. To evaluate pseudo-replication due to the multiple contributions of some mothers, we also conducted a binomial analysis considering the mother's identity as a random effect. This analysis failed to account for the fact that only one male can successfully sire each offspring (see above), but the results were very similar to those found using MDC, strengthening our conclusions.

R_{ped} and R_{QG} were significantly correlated with one another, as were the three estimates of MHC dissimilarity (MHC_{diff} , AA_{diff} and S_{diff}) (Table S1). We, therefore, performed separate analyses with each of these measures to address the questions 'does overall genetic dissimilarity influence reproduction' (two analyses, using R_{ped} and R_{QG}) and 'does MHC dissimilarity influence reproduction' (three analyses, using MHC_{diff} , AA_{diff} and S_{diff}).

Within-male MHC diversity was collinear with other potential explanatory variables (Table S1). We, therefore, addressed the question 'does male genotype influence reproduction?' by including MHC_{male} and AA_{male} in separate analyses, but did not attempt to draw conclusions regarding the relative influence of genetic diversity and male heterozygosity on reproduction.

In each analysis we included the age, dominance status (alpha vs. non-alpha), and IR of the potential sire, as these are known to influence the probability that a male reproduced (Charpentier et al., 2005b). The correlation between age and dominance status was very low ($R^2=0.08$), meaning that the two covariates could be included in the same analyses without problems of collinearity. Measures of within-male MHC diversity were not significantly related to male dominance rank (GENMOD procedure with binomial distribution, some males are included twice because they were both alpha and non-alpha during the study, IR: $n=60$, $X^2=1.25$, $P=0.263$; MHC_{male} : $n=49$, $X^2=0.78$, $P=0.378$; AA_{male} : $n=49$, $X^2=0.83$, $P=0.361$; S_{male} : $n=49$, $X^2=0.148$, $P=0.224$).

We used Akaike's information criteria (AIC) to measure and compare the goodness of fit of statistical models. Where variables significantly influenced reproduction we calculated odds ratios as the exponential function of the conditional logit estimate.

RESULTS

MHC genotyping

We identified 34 different Mandrill sphinx Masp-DRB sequences in 155 individual mandrills (Table S2). Sequences were deposited in GenBank (accession numbers: DQ103715-32, DQ103734-46, EU693911-14). Each individual mandrill possessed 1-7 sequences (those possessing a single sequence were homozygous for that sequence). The seven individuals used for cDNA analysis possessed a total of 16 different *Masp-DRB1*03*, *1*04*, *3*, *5*, **W* and *6* sequences. We identified 15 cDNA *Mhc-DRB* sequences in these individuals, suggesting that most (15/16) of the mandrill MHC-DRB sequences were expressed and possibly functional. The one sequence that was undetected using cDNA had a 1bp deletion, which would disrupt the sequence reading frame and render it incapable of making a functional protein. This sequence was removed from subsequent analyses. Fourteen sequences were assigned to the *Masp-DRB1*03*, *1*04*, *3*, *5* and **W* loci and lineages, representing all loci and lineages known to exist in mandrills. Unexpectedly, the additional expressed sequence (*Masp-DRB6*0403*) was assigned to the *-DRB6* locus, typically a non-functional pseudogene in other primates (Klein & O'hUigin, 1995).

Each nucleotide sequence resulted in a unique amino acid sequence, with the exception of one pair (*Masp-DRB*W301* and *-DRB1*0402*), which differed in nucleotide sequence but encoded the same amino acid sequence. Of 75 amino acid positions, 59 were variable and sequences differed at a mean of 18.3 ± 0.2 sites. Supertype analysis identified 11 MHC-DRB supertypes, containing 1-6 sequences each (Table S2). Of these, S2 was composed only of *-DRB6* sequences. We conducted all subsequent analyses both with and without this supertype, because our cDNA study identified one *DRB6* sequence that appeared to be expressed.

Patterns of reproduction and sample size

The range of variation in values for the various genetic variables investigated is presented in Table S3. The age, rank (alpha vs. not alpha) and IR of a potential sire all significantly influenced which male sired an individual offspring (Table 1). Alpha males

sired 148 offspring (76%), while non-alpha males sired 45 (see also Charpentier et al. 2005). Alpha males were 18 times more likely to sire a given offspring than non-alpha males, older males were more likely to sire than younger males, and male IR was negatively related to the chances of siring, confirming previous results that showed that males with high microsatellite heterozygosity have higher reproductive success in this colony (Charpentier et al., 2005b).

Relatedness and reproduction

The range of values for R_{ped} and R_{QG} are presented in Table S3. R_{ped} significantly influenced the probability of reproduction, which decreased as relatedness increased (Table 1). R_{QG} showed a non-significant trend towards the same effect, but AIC values for the two models were very similar (368 vs. 369, Table 1). Replacing the continuous R_{ped} variables with a cut-off point at $R=0.25$ made very little improvement to the fit of the model (estimate \pm SE: 1.21 ± 0.45 , $t_{179}=2.67$, $P=0.008$, AIC 365).

MHC-dissassortative mating

Mothers possessed 2-7 MHC sequences ($n=34$, mean 3.9 ± 0.2), while potential sires possessed 2-6 sequences ($n=40$, mean 4.0 ± 0.2). Both mothers and potential sires possessed 2-6 MHC supertypes (mothers mean 3.6 ± 0.2 , potential sires mean 3.9 ± 0.1). The range of values for the various measures of MHC dissimilarity in a dyad is presented in Table S3. The probability of reproduction by a given sire increased as MHC_{diff} and AA_{diff} increased (Table 2, Fig 1). In each case the probability of reproduction increased by 17% for each additional MHC sequence or amino acid position that differed (odds ratio 1.17). However, the probability of reproduction did not increase significantly with S_{diff} (Table 2). When we added a quadratic effect of MHC_{diff} to the model we found no significant influence on the probability of reproduction (estimate \pm SE: -0.01 ± 0.02 , $t_{179}=-0.71$, $P=0.48$), suggesting no evidence of choice for intermediate MHC diversity in offspring. AIC was lowest (by a small margin) for the model with AA_{diff} , suggesting that this was the best predictor of reproduction among the MHC variables that we tested.

When we included R and MHC_{diff} in the same model, MHC_{diff} remained a significant influence on reproduction with R_{QG} and $R_{<0.25}$ and showed a tendency to do so with R_{ped} while the influence of R was non-significant in each case (Table 3). Adding R to the

model with MHC_{diff} increased the AIC minimally (Table 3). This suggests that the influence of MHC dissimilarity on reproduction may be stronger than that of overall genetic dissimilarity. However, R and MHC_{diff} were collinear (Table S1), which increases uncertainty in the coefficient estimates. To circumvent this problem, we examined only dyads where $R < 0.25$ (excluding father/daughter dyads and half-siblings) and found that MHC_{diff} was no longer a significant influence on reproduction (Table 3). Nevertheless, this analysis excludes the least MHC-dissimilar dyads, meaning that we cannot distinguish between the two influences definitively. When we included the variable $R < 0.25$ in the same model as MHC_{diff} , only MHC_{diff} was a significant influence on reproduction (Table 3).

Male genotype and reproduction

The range of values for the various measures of within-male MHC diversity is presented in Table 1. MHC_{male} , AA_{male} and S_{male} were not significantly related to IR_{male} (Table S1) suggesting that neutral heterozygosity and adaptive MHC variability were not linked in these males. AA_{male} significantly influenced the probability that a male sired a given offspring (Table 2, Fig. 2) with a 7% increase in the probability of reproduction for each additional amino acid position that differed. However, there was no significant influence of either MHC_{male} or S_{male} on the probability of reproduction (Table 2). This suggests that the amino acid sequence diversity of the MHC genotype of the male was more important in reproduction than the simple number of sequences or supertypes he possessed. However, the P value for AA_{male} was close to 0.05 (0.044), and given that we also tested two other measures of within-male MHC diversity (MHC_{male} and SS_{male}) this may represent a type 1 error. There was no significant influence of the possession of individual supertypes on the probability that a male sired (Table S4).

DISCUSSION

We genotyped a large population of mandrills for MHC-DRB, and demonstrated that many of the MHC sequences we identified via genomic DNA analysis are expressed. Together with previous results showing significantly higher rates of non-synonymous than synonymous substitutions within the mandrill DRB (Abbott et al. 2006), this suggests that the MHC sequences are capable of providing resistance to pathogens, and

thus might be the foundation of MHC-associated mate choice. However, expression is not proof of functionality. For example, although different MHC loci are expressed in the bank vole, only one is under positive selection and associated with parasite resistance, while another expressed MHC locus is not under selection, (Axtner & Sommer, 2007). We are currently investigating the association between specific MHC sequences and parasite resistance in our study population.

The nature of our large dataset, which involves reproduction over multiple years for a long-lived species and collinearity between measures of genetic similarity, poses a problem for statistical analyses. However, using the best statistical models currently available, we found that pedigree relatedness, overall genetic dissimilarity, MHC dissimilarity (number of different MHC sequences and amino acid difference) and male genotype (overall genetic diversity and MHC amino acid diversity) all influenced reproduction in this mandrill colony. The influence of MHC dissimilarity on reproduction appeared to be stronger than that of overall relatedness (R), which was only borderline significant. However, this pattern may still be driven by females simply avoiding brothers/fathers as mates, or low fertilization success if these males do inseminate a female, because when we excluded closely related dyads (who are also least MHC-dissimilar) from our analyses, we found that MHC dissimilarity was no longer significant.

Given the polygynous mating system, strong sexual dimorphism and high male reproductive skew that occur in mandrills, it is quite surprising that other genetic factors also predict which male reproduces. Male rank was by far the strongest influence on reproduction in males, with alpha males being 18 times more likely to sire any given offspring. The nature of our study population limits our power to draw general conclusions on MHC-associated mate choice in wild mandrills, because female choice in our study population is limited to natal males (although these may not related to the female). However, it is interesting to note that findings of MHC-associated mate choice in humans are also from small or isolated populations with little or no migration to introduce genetic variation (Ober *et al.*, 1997; Chaix *et al.*, 2008), situations analogous to the mandrill colony studied here, suggesting that MHC-associated mate choice may be stronger, or easier to detect, under such conditions.

Despite the limitations of the colony environment, our results are broadly similar to those found in previous studies of strepsirrhine primates living in very different social systems: in fat-tailed dwarf lemurs MHC supertype dissimilarity (but not sequence or amino acid dissimilarity) significantly influenced reproduction, and specific superotypes were also associated with male reproductive success (Schwensow et al., 2007a). In grey mouse lemurs sires were more dissimilar to the mother at the level of amino acid sequences, and had more MHC superotypes (but fewer MHC sequences) than randomly assigned males, but no specific superotypes influenced reproduction (Schwensow et al., 2008). In the only other study of MHC-associated mate choice in a non-human anthropoid, male rhesus macaques heterozygous at the MHC-DQB1 locus were found to have greater reproductive success than homozygous males, but MHC-dissimilarity did not influence mate choice (Sauermann et al., 2001). Our results suggest that MHC-associated mate choice may be widespread across the order primates, although the exact patterns observed differ between species. Moreover, our results are the first to demonstrate a reproductive advantage associated with MHC dissimilarity (and possibly MHC diversity measured as amino acid diversity) in a polygynous species with high levels of male-male competition, and suggest that MHC-associated mate choice may be more widespread across different mating systems than previously thought (Paterson & Pemberton, 1997).

Dissassortative mating

Choosing a genetically dissimilar reproductive partner may serve two functions: as a mechanism to avoid inbreeding (Grob *et al.*, 1998; Jordan & Bruford, 1998); or to increase MHC diversity in offspring, improving their ability to recognise and react to a broader range of pathogens, and rendering them fitter than less diverse individuals (Doherty & Zinkernagel, 1975). Our results suggest that the influence of MHC dissimilarity on reproduction was stronger than that of overall genetic dissimilarity, and that mandrills aim to ensure MHC diversity in their offspring. This would result in offspring that were able to respond to a broader range of antigens than less MHC diverse individuals (Doherty & Zinkernagel, 1975). Such pathogen resistance may be particularly important in mandrills, which live in tropical rainforest, and can form very large groups in the wild (Abernethy et al., 2002). Both larger group sizes (Davies et al.,

1991) and wetter environments (McGrew et al., 1989) have been shown to lead to higher rates of parasite infection in primates, and annual rainfall is also positively related to immune system parameters, suggesting that primates living in wetter habitats have evolved to combat a higher risk of disease infection (Semple et al., 2002). Finally, we found no evidence that mandrills choose for an intermediate level of MHC diversity to ensure optimal parasite resistance in their offspring (e.g. Wegner et al. 2003), suggesting that they are choosing for maximum MHC diversity, rather than an intermediate level.

These findings raise the question of how female mandrills select genetically complementary mates. As noted above, we cannot rule out 'standard' inbreeding avoidance of close kin as opposed to finer-scale discrimination among genotypes. Mandrills are female philopatric (Setchell, 1999), and the best indicator of pedigree relatedness of a potential mate may be whether he was born into the same group. However, mandrills live in very large groups in deep rain-forest (Abernethy et al., 2002) and this information may not necessarily be available to females. Moreover, MHC-dissimilarity was a stronger predictor of which male sired a given offspring than pedigree relatedness. MHC-disassortative mating requires comparison of the MHC genotype of potential mates with the chooser's own genotype. Both pre- and post-copulatory mechanisms of female choice may play a role here. Female mandrills are able to express mate choice at the pre-copulatory level (Setchell 2005). The possibility that primates employ self-referent phenotype matching has attracted renewed attention recently (Widdig et al., 2001), and mandrills appear to be able to discriminate paternal kin from non-kin, despite their polygynandrous mating system (Charpentier et al., 2007). The mechanism underlying this phenomenon remains unknown, but it may occur via visual, olfactory, acoustic, or behavioural cues (Widdig, 2007). In this context, it is striking that both male and female mandrills possess a sternal gland which produces a glandular secretion (Feistner, 1991). If genetic similarity at the MHC is reflected in similar odour profiles, then olfaction may play a role in the assessment of mate compatibility, as demonstrated for both rodents and humans (review in Penn 2002).

Female mandrills mate with multiple males during their fertile phase (Setchell, unpublished observations) and genetic compatibility may be determined at the post-

copulatory level via selective fertilisation and/or selective abortion (Zeh & Zeh, 2003; Ziegler *et al.*, 2005). MHC molecules are known to be expressed on the surface of spermatozooids (Paradisi *et al.*, 2000), and mouse oocytes are able to select sperm based on MHC genotype (Wedekind *et al.*, 1996) suggesting that selective fertilisation may potentially account for the observed patterns of reproduction. MHC-associated post-copulatory mate choice has been suggested for grey mouse lemurs, where no difference was found in the MHC genotype of mated and non-mated males in the vicinity of a receptive female, but sires were more dissimilar to the mother at the MHC than randomly assigned males (Schwensow *et al.*, 2008).

Male genotype

Reproduction in the mandrills was also influenced by the genetic characteristics of potential sires, in terms of both neutral (microsatellite) heterozygosity (see also Charpentier *et al.*, 2005b) and MHC amino acid sequence diversity. These results suggest that individual genetic characteristics in mandrills may be linked to male vigour and we are currently investigating whether any or all of microsatellite heterozygosity, MHC diversity, and the possession of particular supertype, are linked to better condition or reduced susceptibility to disease. Higher levels of microsatellite heterozygosity are known to bring general fitness advantages (review in Hansson & Westerberg, 2002), for example via increased metabolic efficiency (Mitton *et al.*, 1993), and this is true for our study population (Charpentier *et al.* 2005, 2006). Increased MHC diversity may also allow a male to resist a greater variety of parasites (review in Penn *et al.* 2002). These results may thus reflect intrasexual competition, with MHC diversity conferring superior competitive ability on particular males. However, our analyses included the influence of male dominance rank as a separate variable, implying that intersexual selection (mate choice) may also be occurring for genetic characteristics.

Males are unable to pass on heterozygosity at specific loci (Brown, 1997; Mays & Hill, 2004) and heterozygous males have therefore been thought to confer direct, rather than indirect, fitness benefits on their offspring (Partridge, 1983). However, heterozygous males also sire offspring that are themselves more heterozygous, on average (Mitton *et al.*, 1993), and possess more rare alleles than homozygotes, which can be inherited by offspring (Apanius *et al.*, 1997), suggesting that females may also receive indirect

benefits from genetically diverse mates. Indeed, a recent theoretical model has shown that directional mating preferences for heterozygous males can evolve and be maintained in the absence of direct fitness benefits (Fromhage *et al.*, 2009). Genome-wide heterozygosity has been suggested to act as a marker of MHC diversity (Aparicio *et al.*, 2001; Acevedo-Whitehouse *et al.*, 2003), such that mate choice for males signaling general genetic diversity leads to choice for MHC-diverse mates. However, the reverse has also been suggested: that MHC diversity may act as a marker of genome-wide heterozygosity (Penn & Potts, 1999). In this context, it is interesting that we found no significant relationship between neutral heterozygosity and adaptive MHC diversity in males, suggesting that the two measures of diversity are independent in mandrills, and that females would be unable to use one as a marker for the other.

If the increased reproductive success enjoyed by genetically diverse males is due to increased vigour in these males, which is preferred by females, then this raises the question of how heterozygosity and MHC diversity are signaled to females. Male mandrills possess a suite of secondary sexual ornaments, including bright red coloration on the face, rump and genitalia (Hill, 1970), and females prefer to mate with redder males (Setchell, 2005). If sexually selected traits signal the possession of 'good genes' (Zahavi, 1975; Hamilton & Zuk, 1982; Brown, 1997) in mandrills, in the form of genome-wide or MHC diversity, then males with more exaggerated ornaments should also be genetically more diverse. Condition dependent secondary sexual traits have been shown to correlate positively with overall genetic diversity in a variety of other species, including birds (Aparicio *et al.*, 2001; Foerster *et al.*, 2003; Marshall *et al.*, 2003), fish (Müller & Ward, 1995; Sheridan & Pomiankowski, 1997; van Oosterhout *et al.*, 2003), and invertebrates (Aspi, 2000). At the level of the MHC, male pheasants with particular MHC genotypes have larger spurs, a trait which is known to influence female choice (von Schantz *et al.*, 1997). Certain MHC genotypes are also associated with antler size and body size in white-tailed deer (*Odocoileus virginianus*) (Ditchkoff *et al.*, 2001). Antler size in this species is also related to helminth abundance, suggesting that antlers honestly advertise the possession of good genes for parasite resistance (Ditchkoff *et al.*, 2001).

In conclusion, we demonstrate both sexual selection for genetic complementarity (MHC dissimilarity) and directional selection for good genes (genome-wide heterozygosity and MHC diversity) in a primate species living in large multi-male, multi-female groups. This implies that female mandrills employ a combination of mate choice strategies, as the male with the best genes may not be the most genetically compatible mate for every female. For example, they may switch between the two mate choice strategies according to the available diversity of males (Roberts & Gosling 2003), or employ a hierarchical, nested model of mate choice, in which they choose the most compatible male from the subset of males possessing good genes (Mays & Hill, 2004). Our results are the first to demonstrate mate choice for genetic dissimilarity in a species characterised by high reproductive skew among males, and suggest that MHC-associated mate choice can occur even where male-male competition is intense. Finally, our results concern an isolated population with no migration to introduce genetic variation, a situation analogous to those in which MHC-associated mate choice has been found in humans, suggesting that MHC-associated mate choice may be especially important in such populations.

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882

Table 1: The influence of male age, rank, and relatedness to the mother on the

884 probability that a given male sired an offspring.

Analysis	Parameter	df	Estimate \pm SE	t value (t_{192})	Approx. Pr > t
1	Age	1	1.45 \pm 0.28	5.10	<0.0001
AIC=368	Age ²	1	-0.06 \pm 0.01	-4.61	<0.0001
	Rank	1	2.56 \pm 0.20	12.93	<0.0001
	R _{ped}	1	-1.72 \pm 0.85	-2.02	0.043
	IR	1	-3.85 \pm 0.87	-4.40	<0.0001
2	Age	1	1.42 \pm 0.28	5.01	<0.0001
AIC=369	Age ²	1	-0.06 \pm 0.01	-4.52	<0.0001
	Rank	1	2.57 \pm 0.20	12.98	<0.0001
	R _{QG}	1	-1.05 \pm 0.58	-1.82	0.069
	IR	1	-3.37 \pm 0.88	-3.83	0.0001

886 Results of MDC Procedure, Conditional Logit Estimates.

Analyses were conducted separately for R_{ped} and R_{QG} due to collinearity.

888

Table 2: The influence of MHC dissimilarity and male genotype on the probability that a given male sired an offspring.

Analysis	Parameter	df	Estimate \pm SE	t value (t_{179})	Approx. Pr > t
1	MHC _{diff}	1	0.17 \pm 0.07	2.33	0.020
AIC=301	MHC _{male}	1	-0.10 \pm 0.15	-0.70	0.49
	Age	1	1.56 \pm 0.41	3.85	0.0001
	Age ²	1	-0.06 \pm 0.02	-3.69	0.0002
	Alpha (0/1)	1	2.64 \pm 0.23	11.26	<0.0001
	IR	1	-3.90 \pm 1.10	-3.53	<0.0001
2	AA _{diff}	1	0.16 \pm 0.06	2.67	0.008
AIC=298	Age	1	1.60 \pm 0.42	3.85	0.0001
	Age ²	1	-0.06 \pm 0.02	-3.66	0.0002
	Alpha (0/1)	1	2.66 \pm 0.23	11.34	<0.0001
	IR	1	-3.67 \pm 1.04	-3.53	0.0004
3	AA _{male}	1	0.07 \pm 0.03	2.01	0.044
AIC=302	Age	1	1.60 \pm 0.41	3.88	0.0001
	Age ²	1	-0.06 \pm 0.02	-3.70	0.0002
	Alpha (0/1)	1	2.65 \pm 0.23	11.52	<0.0001
	IR	1	-3.28 \pm 1.04	-3.14	0.002
4	S _{diff}	1	0.10 \pm 0.08	1.24	0.22
AIC=306	S _{male}	1	0.07 \pm 0.17	0.39	0.70
	Age	1	1.51 \pm 0.41	3.71	0.0002
	Age ²	1	-0.06 \pm 0.02	-3.49	0.0005
	Alpha (0/1)	1	2.63 \pm 0.22	11.70	<0.0001
	IR	1	-3.98 \pm 1.08	-3.68	0.0002

Results of MDC Procedure, Conditional Logit Estimates

Analyses were conducted separately due to collinearity of the different estimates of MHC dissimilarity.

Conducting supertype analyses without S2 (because S2 comprised only -DRB6 sequences, which may be non-functional) did not alter the significance of our results.

898 Table 3: Comparing the influence of overall genetic dissimilarity and MHC-dissimilarity
on the probability that a given male sired an offspring.

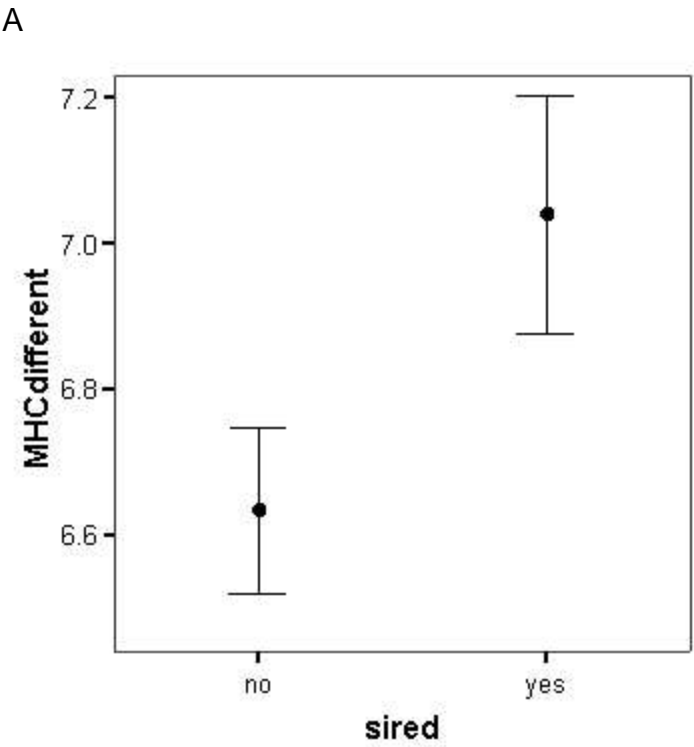
Analysis	Parameter	df	Estimate \pm SE	t value (t_{179})	Approx. Pr > t
1	R _{ped}	1	-0.83 \pm 1.06	-0.78	0.43
AIC=303	MHC _{diff}	1	0.12 \pm 0.07	1.70	0.09
	Age	1	1.57 \pm 0.41	3.83	0.0001
	Age ²	1	-0.06 \pm 0.02	-3.67	0.0002
	Rank	1	2.66 \pm 0.23	11.53	<0.0001
	IR	1	-3.72 \pm 1.05	-3.55	0.0004
2	R _{QG}	1	-0.35 \pm 0.68	-0.52	0.60
AIC=302	MHC _{diff}	1	0.13 \pm 0.06	2.11	0.03
	Age	1	1.54 \pm 0.40	3.80	0.0001
	Age ²	1	-0.06 \pm 0.02	-3.64	0.0003
	Rank	1	2.67 \pm 0.23	11.58	<0.0001
	IR	1	-3.58 \pm 1.06	-3.39	0.0007
2	R _{<>0.25}	1	0.13 \pm 0.40	0.32	0.748
AIC=302	MHC _{diff}	1	0.15 \pm 0.07	2.30	0.021
	Age	1	1.54 \pm 0.40	3.81	0.0001
	Age ²	1	-0.06 \pm 0.017	-3.64	0.0003
	Rank	1	2.68 \pm 0.23	11.54	<0.0001
	IR	1	-3.63 \pm 1.05	-3.46	0.0005

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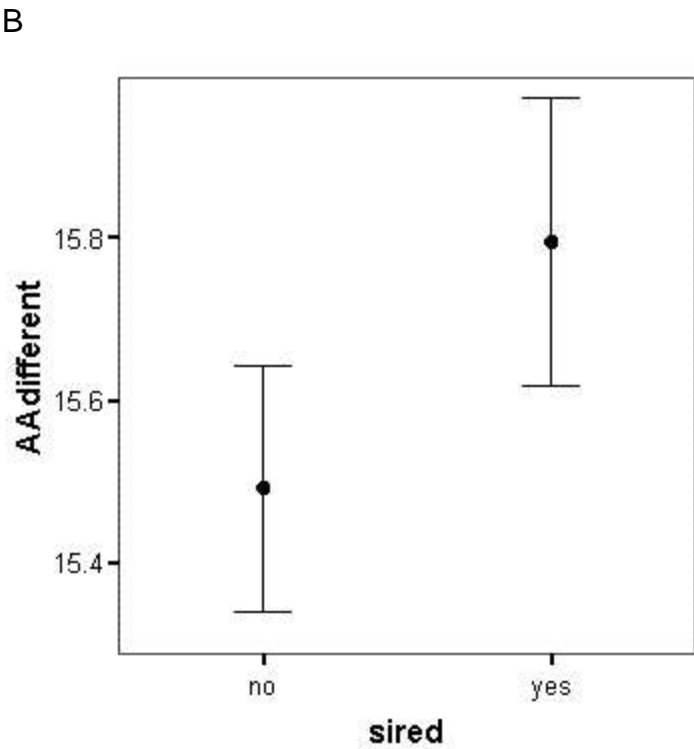
Results of MDC Procedure, Conditional Logit Estimates

902 Analyses were conducted separately for R_{ped} and R_{QG} due to collinearity.

904 Fig. 1. Influence of MHC dissimilarity on whether reproduction occurred. Figure
compares mean \pm se MHC_{diff} (A) and AA_{diff} (B) for the sire of each offspring with the
906 mean value for non-sires for each individual offspring (n=180 offspring).



908



910

Fig. 2. Influence of AA_{male} on whether reproduction occurred. Figure compares the mean \pm sem AA_{male} for the sire of each offspring with the mean value for all the non-sires of that offspring (n=180 offspring).

